# ON THE MECHANISM OF HETEROGENEOUS LIQUID-PHASE OXIDATION OF CUMENE, CYCLOHEXENE AND TETRALIN

A Thesis Submitted
In Partial Fulfilment of the Requirements
for the Degree of
MASTER OF TECHNOLOGY

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to the
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# CERTIFICATE

Certified that this work has been carried out under our supervision and it has not been submitted elsewhere for a degree.

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### 1. INTRODUCTION

Recently, the heterogeneous catalytic oxidation of arenes namely, cumene and tetralin and the cyclic compound, cyclohexene, have been subjects of numerous investigations [1-10]. The mechanisms of these reactions have drawn considerable academic and industrial attention. In the past five years there has been considerable advancement in the elucidation of the mechanisms of oxidations of cumene and cyclohexene. A brief literature survey on the mechanisms of oxidation of these compounds is presented in the following section:

# 1.1 Literature Survey:

Mukherjee and Graydon [1] have studied the liquidphase oxidation of tetralin (1,2,3,4-tetrahydronaphthalene)
using both soluble and insoluble catalysts and compared the
reaction rates. Oxides of nickel, manganese, and copper were
found to be extremely active while those of aluminium and
zinc were inactive. Rates of oxidation were measured in the
temperature range of 45-90°C. The initial product of oxidation
reaction was found to be tetralin hydroperoxide which decomposed
further into tetralone and tetralol. A ketone:alcohol ratio
of 2:1 was observed in the product with most of the catalysts.
It was further noticed that a critical hydroperoxide to
catalyst ratio existed below which the reaction did not proceed.

Kinetic studies have been made with four of the best catalysts, and the reaction mechanism proposed is as follows:

Initiation 
$$\frac{k_1}{ROOH \cdot M} \xrightarrow{k_2} RO \cdot M + OHM$$
 $RO \cdot M + RH \xrightarrow{k_3} ROH + R$ 
 $OHM + RH \xrightarrow{k_3} H_2O + R$ 

Propagation

$$R^{\bullet}+O_{2} \xrightarrow{k_{4}} RO_{2}^{\bullet}$$

$$RO_{2}^{\bullet} + RH \xrightarrow{k_{5}} ROOH + R^{\bullet}$$

Termination (bimolecular)

$$2 \text{ RO}_{2}^{*} \xrightarrow{k_{6}} 2R \text{ " CO} + H_{2}\text{O} + O_{2}$$

Termination (monomolecular)

$$RO_{2}^{\bullet} \xrightarrow{k_{7}} R''CO + O_{2} + H_{2}O$$

$$RO_{2}^{\bullet} \xrightarrow{k_{8}} R''CO + M'OH$$

with  $R = C_6H_5C(CH_3)_2$  and M = active site of the catalyst surface

Van Ham et al. [2] have found that the use of silver catalyst showed a significant lowering of the activation energy in the oxidation of cumene. They also found a pronounced increase in the oxidation rate by using silver instead of silver-gold alloy as the catalyst.

Neuburg et al. [3] studied the heterogeneous liquidphase oxidation of cyclohexene using manganese dioxide as the catalyst. The decomposition of cyclohexenyl hydroperoxide was also studied. The dependence of initial rate on the catalyst emount had been studied both for exidation and decomposition reactions and a degenerate chain branching mechanism had been postulated by the authors. Hydroperoxide which was produced in the propagation step was decomposed on the solid generating the free radicals necessary for the initiation of the chains. Cyclohexenol and cyclohexenone were produced in the initiation and termination steps. Above a critical catalyst to hydrocarbon ratio, no exidation reaction was observed. At that critical composition, the rate of hydroperoxide decomposition was equal to its rate of formation. The length of the chain decreased to a low value with increasing catalyst to hydrocarbon ratios. The mechanism adapted is as follows:

Initiation

ROOH + M 
$$\xrightarrow{k_1}$$
 RO' + M. HO'

RH + M. HO'  $\xrightarrow{k_2}$  R'+M+H<sub>2</sub>O

RO' + RH  $\xrightarrow{k_3}$  ROH + R'

Propugation

$$R^{\bullet} + O_{2} \xrightarrow{k_{4}} RO_{2}^{\bullet}$$

$$RO_{2}^{\bullet} + RH \xrightarrow{k_{5}} ROOH + R^{\bullet}$$

Homogeneous termination

$$RO_{2}^{2} + RO_{2}^{2} \xrightarrow{k_{6}} R' = 0 + ROH + O_{2}$$

$$RO^{2} + RO^{3} \xrightarrow{k_{7}} R' = 0 + ROH$$

The oxidation of cumene and the decomposition of cumene hydroperoxide on silver, copper and platinum were studied by Casemier et al. [4]. Cumene hydroperoxide dissolved in dodecane

rapidly incomposed by platinum and appreciably by silica geler by silver in the absence of gaseous oxygen, but virtually no decomposition was observed with unsupported silver in an oxygen atmosphere. The oxidation of cumene to cumene hydroperoxide showed a completely different picture; this oxidation was strengly catalyzed by silver, whereas copper, gold and platinum did not catalyze the reaction. For the cumene oxidation with silver a free radical (presumably R00°) was detected in the reaction mixture by EPR spectroscopy. The reaction mechanism assumed diatomic oxygen to be adsorbed on the silver surface. The proposed mechanism is as follows:

Initiation

Property tion

$$R^{\bullet}O_2 + RH \xrightarrow{fast} ROO^{\bullet}$$
 $ROO^{\bullet} + RH \xrightarrow{} ROOH + R^{\bullet}$ 

Termination

2 ROO' ---- non radical product.

Verme and Graydon [5] studied the heterogeneous catalytic oxidation of cumene in liquid phase. The investigation was made at 80°C with molecular oxygen. Cobalt II, III oxide, MnO<sub>2</sub>, NiO and Cu<sub>2</sub>O were used as catalysts. Detailed studies of the effect of catalyst weight, hydrocarbon concentration and initial

hydrop roxide concentration on the rate of oxidation were conducted employing cobult II, III oxide as the catalyst. The expense absorption rate was found to be 0.4 power with respect to the catalyst weight, and 1.5 power with respect to the camera consentration. The concentrations of the main products of exidation of comenc, viz., cumene hydroperoxide, dimethyl charged carrinol and acetophenone were found to depend on the weight of the catalyst. Reaction mechanisms and rate expressions for the exidation of cumene were proposed. The mechanisms are illustrated below:

Initiation
$$RH + M \xrightarrow{k_1} R^*$$
Proposition
$$R^* + C_2 \xrightarrow{k_2} ROO^*$$

$$RCO^* + RH \xrightarrow{3} ROOH + R^*$$

Termination  $\frac{k_4}{R00^{\circ} + R00^{\circ}}$  inactive products

The above mechanism was postulated when there was no hydroperoxide present in cumene. The effect of the initial hydroperoxide would be felt in the initiation reaction and a possible mechanism proposed was:

ROOH...M 
$$\frac{k_5'}{}$$
 RO\*...M + \*OH...M,  
RO\*...M + RH  $\frac{k_6}{}$  ROH + R\*  
\*OH...M + RH  $\frac{k_7}{}$  H<sub>2</sub>O + R\*

Neuburg et al. [6,7] conducted a detailed investigation on the oxidation of cyclohexene and decomposition of cyclohexene hydroperoxide. The liquid-phase decomposition as also the oxidation of cyclohexene hydroperoxide in cyclohexene [6] had been studied in the temperature range of 40 - 70°C using three types of MnO, with different specific surfaces as The influence of initial hydroperoxide concentration catalysts. catalyst weight to liquid volume ratio and temperature were studied in each case. It was postulated that the hydroperoxide formed an equilibrium complex with the catalyst which further decomposed to yield free radicals. The initial activity of oxides decreased during the reaction and the observed decay in the rate of hydroperoxide decomposition had been attributed mainly to the deactivation of catalyst surface. The decomposition products were cyclohexenol, cyclohexenone and water. The number of moles of alcohol formed was higher than the number of moles of ketone in each case. In the oxidation study, reaction periods upto 60 min. were used to study the initial activity of the catalyst. The maximum rate of cyclohexene oxidation was 0.5 order with respect to the catalyst in the region of low catalyst weight to hydrocarbon ratios. A much lower order was found for higher ratios. critical amount of catalyst, the oxidation was completely inhibited due to the quantitative decomposition of hydroperoxide. The maximum oxidation rate was proved to be first order with

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respect to the concentration of cyclohexene. The oxidation rate was of zero order with respect to the initial hydroperoxide concentration in the range of 3 x  $10^{-3}$  to 0.5gmole/liter. giving support to the theory that free radicals were produced from the slow decomposition of the catalyst-hydroperoxide Oxidation products such as cyclohexenol and cyclohexenone inhibited the oxidation rate when they were initially present in concentrations greater than 50 µmole/ml. activation energies for the oxidation ranged between 11.4 and 13.0 Kcal/mole. Product distribution studies showed that cyclohexenol and cycloxenone were produced in nearly equal amounts. A degenerating chain branching mechanism was proposed in which the catalyst played an important role in the initiation and termination stops. Rate equations derived from this mechanism ex lained quantitatively the experimental observations.

Panayotova et al. [9] worked on the role of metallic copper in the exidation of cumene to cumene hydroperoxide.

They reported that at moderate temperatures copper protects cumene hydroperoxide against decomposition. Copper corrosion takes place upon continuous processing of raw material, particularly at high temperatures. Interaction between the metal extracted into reaction medium and cumene hydroperoxide lead to intense decomposition of cumene hydroperoxide.

Srivastava and Srivastava [8] investigation the liquid phase oxidation of cumene with  $\mathrm{Cr}_2\mathrm{O}_3$ ,  $\mathrm{MnO}_2$  and  $\mathrm{Fe}_2\mathrm{O}_3$  catalysts

in the temperature range of 70 - 90°C. Similar to the earlier finding of Varma and Graydon, the present authors observed a critical hydroperoxide to catalyst ratio, below which the reaction did not proceed. Effects of catalyst weight and hydrocarbon concentration on the rate of oxidation were studied in detail. The apparent activation energy for the overall oxidation was calculated to be 9.8 Kcal/g mole. A reaction mechanism had been proposed and the rate expression derived was in good agreement with the experimental data.

Agarwal and Srivastava [10] studied the kinetics and catalytic behaviour of NiO supported on alumina and activated carbon and also Ni MoO, supported on alumina in the liquid phase oxidation of cumene. The catalysts had been characterized by means of DTA, TGA, X-ray, IR and BET studies. Detailed studies of the concentrations of main products along with effect of catalyst weight, and hydrocarbon concentration on the rate of oxidation were conducted at 80°C employing these supported catalysts. A reaction mechanism was proposed in which the catalyst plays an important role in the initiation and termination steps. Rate equations derived were in good agreement with the experimental data. It was observed that use of carbon as a support instead of alumina (NiO catalysts) induced increased activity, lower activation energies, and shifted the limiting oxidation rate to lower catalyst amount.

It has been established that, for the heterogeneous liquid phase oxidation of cumene, cyclohexene and tetralin catalyzed by transition metal oxides, there is a critical weight of catalyst above which the rate of oxidation drops catastrophically [1,3,5,8], which suggested that the rates of formation and destruction of hydroperoxide became equal [7]. A general mechanism explaining the simultaneous formation and destruction of hydroperoxide in the oxidation of cyclohexene has recently been proposed by Neuburg et al. [7], and used by Agarwal and Srivastava [10] to explain the kinetics of cumene oxidation. However, very few data have been presented for these systems in the region where inhibition was effective.

#### 1.2 Present Work:

In the present work MnO<sub>2</sub> catalyzed exidations of cumene, cyclohexene and tetralin in the region where inhibition was effective, have been studied at the temperatures of 80, 60, and 65°C respectively. The results favor the view that these catalyzed exidations involve simultaneous formation and destruction of hydroperoxide. Also the chain branched character of the generation of free radicals is confirmed.

#### 2. EXPERIMENTAL

#### 2.1 Materials:

Cumene, cyclohexene and tetralin were obtained from Herdillia Chemicals, Koch-Light (England) and Riedel De Haen Ag (Germany), respectively. Cumene and cyclohexene were further distilled fractionally.

Monochlorobenzene of Sarabhai M. Chemicals was used as inert solvent for varying the concentration of hydrocarbons.

Hydroperoxides of cumene, cyclohexene and tetralin were prepared separately by the thermal oxidation of the respective hydrocarbons at  $60^{\circ}\text{C}$  using MnO<sub>2</sub> as the catalyst.

Oxygen and nitrogen were supplied by Indian Oxygen
Limited, Kanpur. Nitrogen was used as an inert for the decomposition runs.

 ${\rm MnO}_2$  catalyst was supplied by BDH (India) Limited, Bombay.

# 2.2 Apparatus:

The apparatus was essentially similar to that used by Bolland [11] for the thermal oxidation of ethyl linoleate.

The apparatus consisted of a graduated gas-burette.

The burette was 50 cm long and 1 cm inner diameter with both ends open. The bottom was connected to a mercury reservoir through a polythene tubing as shown in Figure 1.

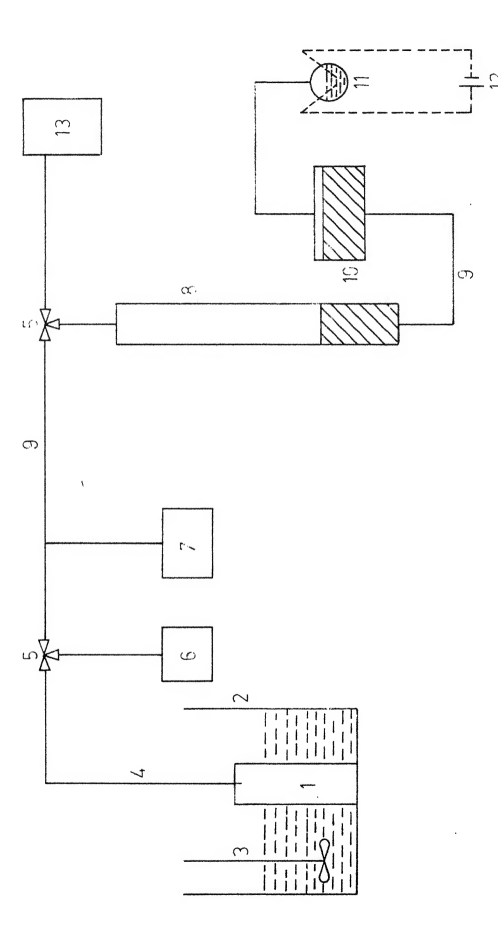


Fig. 1 - Schematic diagram of experimental set-up.

#### KEY TO FIGURE 1

- 1 : B 55/54 Glass Joint Reactor
- 2 : Constant Temperature Water Bath
- 3 : Remi Stirrer
- 4 : Rubber Tubing
- 5 : Three-way Stop Cock
- 6 : Connection to Vacuum Pump through Trap
- 7 : Mercury Manometer
- 8 : Gas-Burette With Mercury
- 9 : Polythene Tubing
- 10 : Mercury Reservoir
- ll : Oxalic Acid Container
- 12 : 12V Battery
- 13 : Connection to Gas Cylinder

Oxygen line and the reactor were connected to the gas-burette by polythene tubing through a three-way stop-cock. The reactor consisted of a B 55/54 glass joint. The male joint was connected to a three-way stop-cock, which in turn was connected to a vacuum line and mercury manometer. The reactor volume was 400 cc. A vacuum pump was used for evacuating the system. The reactor was placed in a constant temperature water bath. Water in the bath was stirred by a femi-stirrer. The bath was heated by a 2 KW heater. An on-off relay and a contact thermometer of range 0-300°C were used to maintain the desired bath temperature.

Efficient stirring of the reaction mixture was achieved by using a magnetic stirrer. The stirrer consisted of a glass-covered iron piece.

# 2.3 Procedure:

All the exidation runs were performed using MnO<sub>2</sub> as the catalyst at the corresponding temperatures mentioned below for different systems.

Cumene 80°C

Cyclohexene 60°C

Tetralin 65°C

The experimental procedure consisted of the following steps:

- 1. The water-bath was switched on and allowed to reach the desired steady-state temperature, which normally took half an hour.
- 2. The desired catalyst amount was taken in the clean and dry reactor. Then the iron stirring piece was dropped in the reactor.
  - 3. 25 ml of the hydrocarbon were added in the reactor.
- 4. The reactor was connected to the gas-burette and the system was evacuated by means of a vacuum pump.
  - 5. The system was filled with oxygen.
  - 6. The reactor was placed in the bath.
- 7. After fifteen minutes the magnetic stirrer was switched on and change in the burette reading was recorded.
- 8. After the run the products were allowed to cool down to room temperature and then centrifuged for fifteen minutes at 800 RPM in order to separate the catalyst from the products.

# Hydroperoxide Decomposition Runs:

For the decomposition runs, the hydroperoxide solution of the respective hydrocarbon of known concentration was added, together with the desired amount of MnO<sub>2</sub> catalyst to the reactor, under nitrogen. All the decomposition runs were performed for thirty minutes.

The samples were analyzed as described below:

#### 2.4 Analytical

The main products of decomposition were the respective hydroperoxide, alcohol and ketone. In the case of cumene, cumene hydroperoxide, cumyl alcohol and acetophenone were the products. Similarly for the decomposition of cyclohexene, cyclohexene hydroperoxide, cyclohexenol and cyclohexenone were the products, while tetralin hydroperoxide, tetralol and tetralone were the products in the decomposition of tetralin. The following methods were used for analyzing the various products of the decomposition runs.

#### hydroperoxide:

The hydroperoxide content in each case was estimated by iodometric method [12]. 40 ml of isopropyl alcohol and 5 ml sample were mixed and refluxed. The mixture was cooled to room temperature and 10 ml isopropyl alcohol saturated with sodium iodide were added and refluxed. The liberated iodide was titrated against sodium thiosumphate solution.

# Alcohol:

Cumyl alcohol and peroxide were converted to phenol by treating the sample with sulfuric acid and hydrogen peroxide[13]. Bromide-bromate method was used to estimate phenol [17]. 5 ml of the sample was treated with 50 ml of 0.lN of potassium bromate-bromide solution and 5 ml of HCL. The mixture was shaken and allowed to stand for 15 min. Two grams of potassium iodide were added. The liberated iodine was titrated with 0.lN sodium thiosulphate using 2 ml of starch indicator.

# Ketone:

The ketone concentration in the product was determined using Perkin-Elmer-137 IR spectrophotometer by studying the absorption at 1700  $\,\mathrm{cm}^{-1}$ .

#### 3. RESULTS AND DISCUSSION

The symbols RH, ROOH and ROH are used in general for hydrocarbons, hydroperoxides and alcohols, respectively. All the relevant experimental data for the oxidation and decomposition runs are presented in Appendices A, B, and C respectively.

In order to study the effect of hydrocarbon concentration in the region where inhibition was effective, it was necessary to study the oxidation rate as a function of the catalyst ratio for all the three systems. The catalyst ratio is defined as the weight of the catalyst in grams divided by the volume of the hydrocarbon in ml.

#### 3.1 Initiation:

The reaction did not commence when the cumene oxidation was performed at  $80^{\circ}\text{C}$  in absence of the catalyst or any free r radical initiator. When some hydroperoxide was added to the system the oxygen consumption started immediately. Therefore, to initiate the reaction both catalyst and hydroperoxide were necessary in all cases. Hydroperoxide concentration to catalyst ratio of approximately 2.5 x  $10^{-4}$  gmole/lit g was used in all the three cases.

# 3.2 Effect of Catalyst Weight:

# Oxidation of Cumene:

The amount of oxygen absorbed versus time of oxidation for cumene at  $80^{\circ}\text{C}$  is shown in Figure 2. Data were obtained

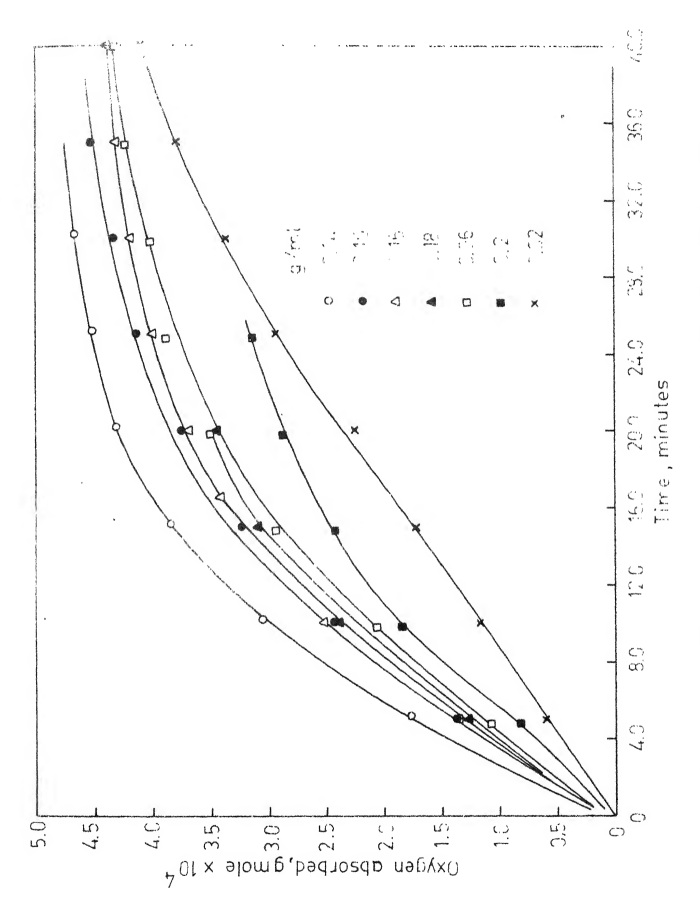


Fig. 2 -Oxygen absorption curves for cumene at 80°C for different MnO2 catalyst weights perform of comene.

on the rate of oxygen absorption for various catalyst weight to hydrocarbon volume ratios ranging from 0.02 to 0.2 gram of catalyst per ml. of hydrocarbon. The initial rate obtained for 0.02 g/ml was 1.7 x  $10^{-7}$  g.mole/sec. Srivastava and Srivastava [8] observed an oxygen absorption rate of 1.3 x  $10^{-7}$  g mole/sec under the same conditions.

The rate of oxygen absorption versus catalyst weight is shown in Figure 3. It was observed that the rate of oxygen absorption was maximum at 0.14 g/ml. A further increase in the catalyst weight resulted in the decrease of rate of oxygen absorbed. This behaviour was observed by Srivastava and Srivastava [8] as well as by Varma and Graydon [5]. The value above which the rate declines is referred to as the critical catalyst amount. Srivastava and Srivastava [8] established that the critical catalyst amount was unaffected by variation in temperature over the temperature range 70 - 80°C.

# Oxidation of Cyclohexene:

The data recorded in Table B.1 of Appendix B represent the consumption of oxygen with time for various catalyst ratios with MnO<sub>2</sub> as a catalyst at 60°C. A detailed study was performed to obtain the rate of oxygen absorbed for various catalyst ratios ranging from 0.08 to 0.4 g/ml. Figure 4 shows the rate against catalyst ratio. As in the case of cumene the rate of oxidation dropped rapidly after 0.2 g/ml. Similar observation was made by Neuburg et al. [3]. They observed a

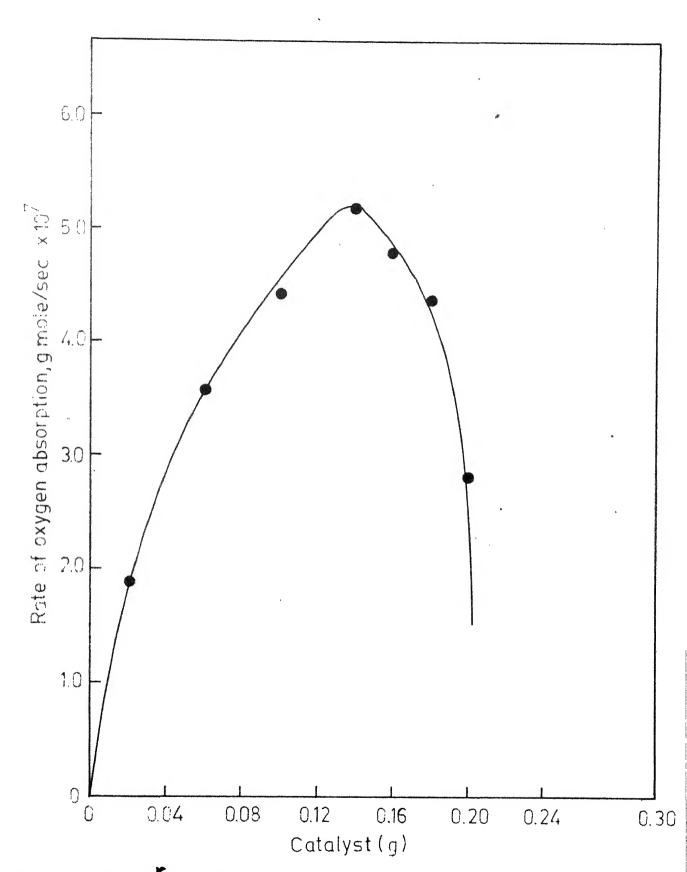


Fig. 3 - Measued rates of oxygen absorption for 1ml. of cumene at 80°C as a function of weight of MnO2 catalyst.

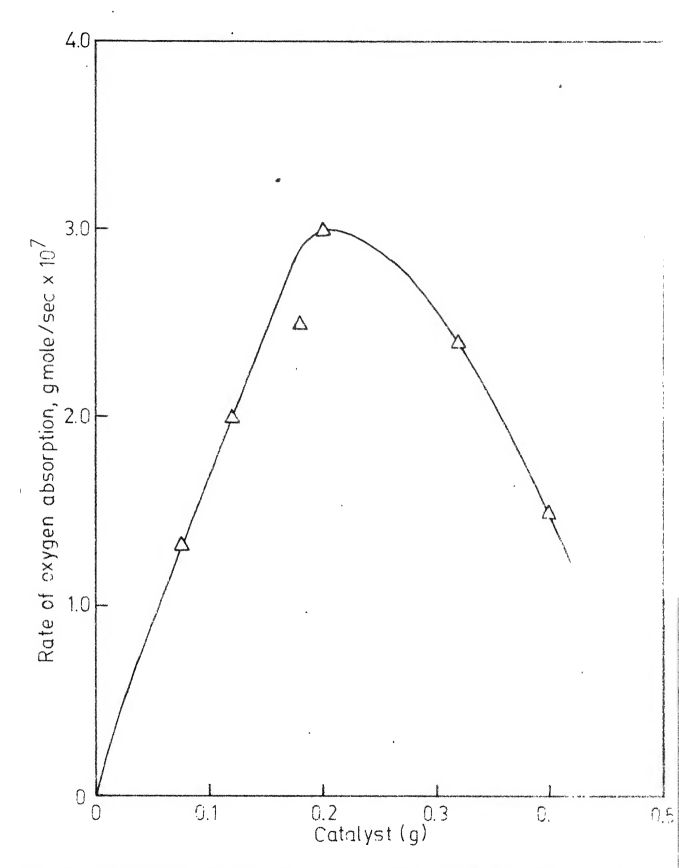


Fig. 4 - Measured rates of oxygen absorption for 1ml cycloheken at 60°C as a function of MnO2 catalyst weight.

decline in rate at 0.16 g/ml under the same conditions. In the present study the maximum rate observed was  $3 \times 10^{-7}$  g.mole/sec. while Neuburg et al. [3] found  $7 \times 10^{-7}$  g.mole/sec.

#### Oxidation of Tetralin:

The oxidation of tetralin was carried out using the same MnO<sub>2</sub> catalyst at 65°C. The amount of oxygen absorbed versus time is shown in Table C-l (Appendix C). Oxygen absorption was studied at catalyst ratios of 0.2, 0.4, 0.8, 0.9 and 1.0 g/ml. The rate data were plotted against catalyst ratios as shown in Figure 5. As seen the behaviour is similar to those of oxidation of cumene and cyclohexene. Mukherjee and Graydon [1] also observed the same behaviour while investigating this reaction.

Thus, in all the three cases of oxidation of cumene, cyclohexene, and tetralin there is a catalyst ratio above which oxidation rates decline. It has already been established that before the break-point the rates of oxidation in the presence of transition metal oxides proceed with approximately 0.5 and 1 power with respect to catalyst ratio and hydrocarbon concentration, respectively.[2,5,8,10].

# 3.3 Effect of Hydrocarbon Concentration in the Inhibiting Region:

The influence of cumene, cyclohexene and tetralin concentrations on the respective rate of oxidation was studied with the mixture of the corresponding hydrocarbon with

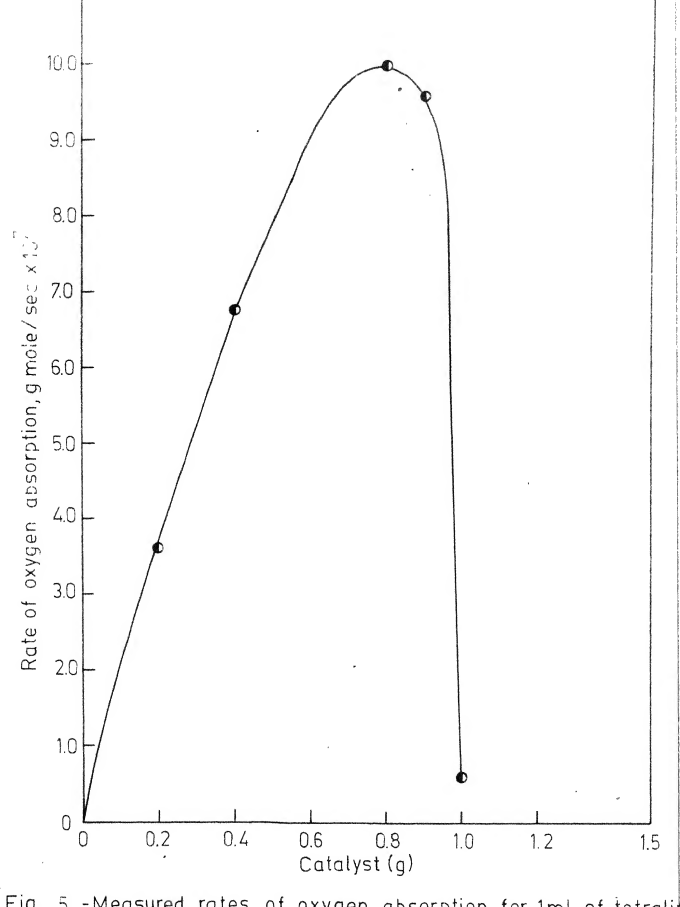


Fig. 5 -Measured rates of oxygen absorption for 1ml. of tetralinat 65℃ as a function of MnO2 catalyst weight.

monochlorobenzene at catalyst ratios chosen from the inhibitir region.

#### Cumene:

Runs were conducted for three different catalyst ratios 0.16, 0.18 and 0.2 g/ml (Figure 3). Figure 6 represents a typical plot of oxygen absorbed versus time at various hydrocarbon concentrations for the catalyst ratio 0.16 g/ml. The data for 0.18 and 0.2 g/ml. are presented in Table A-2 (Append A). The rates were calculated and plotted against the cumene concentration, on a log-log graph as shown in Figure 7. It is interesting to note that the orders obtained were 2.3, 2.1, and 2.2 with respect to hydrocarbon concentration for the catalyst ratios 0.16, 0.18 and 0.2 g/ml, respectively. It shows that the rate dependence with respect to the cumene concentration in the inhibiting region is approximately 2.

# Cyclohexene:

As in the case of cumene the cyclohexene concentration was varied and runs were conducted for four different concentrations. The catalyst ratio chosen was 0.22 g/ml (Figure 4). Table B.2 (Appendix B) represents the data of oxygen absorbed versus time. The rate of oxidation as a function of cyclohexene concentration is shown in Figure 8 on a log-log plot. The rate dependence with respect to cyclohexene concentration was found to be 1.7.

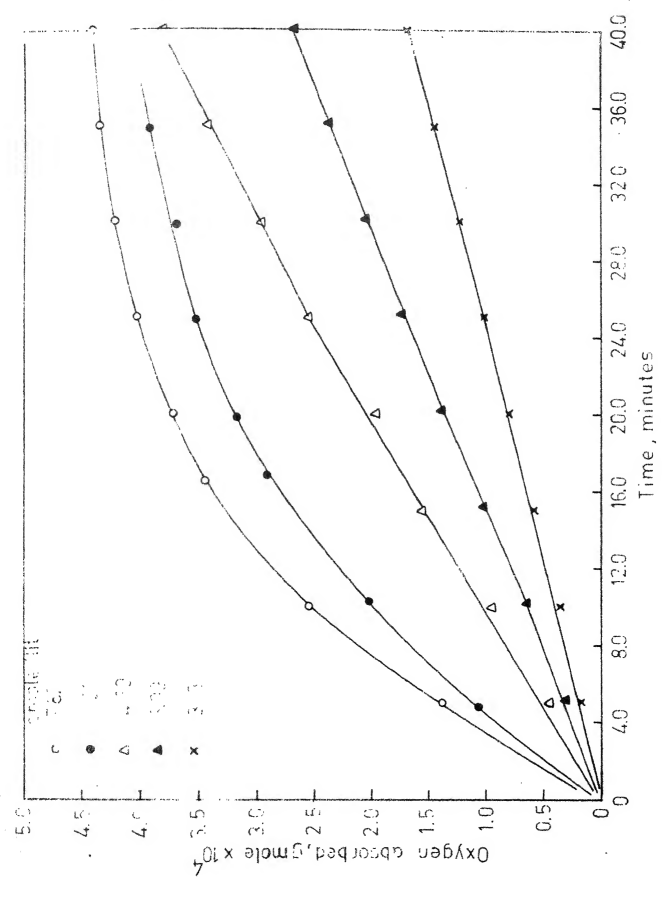


Fig. 5 -Oxygen absorption curves for catalyst ratio 0.15g MnO2 per ml. of cumene at 180°C as a function of cumene concentration(gmole/lit.)

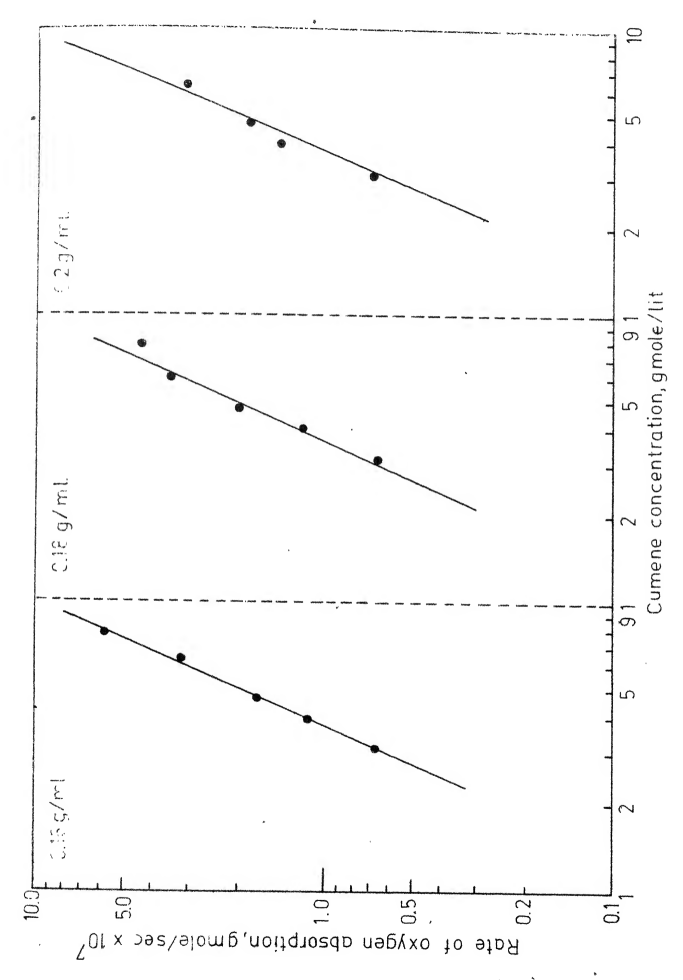


Fig. 7 -Rate dependence with respect to cumene concentration for catalyst weights 0.16g, 0.18g & 0.2g per 1ml of cumene at 83°C

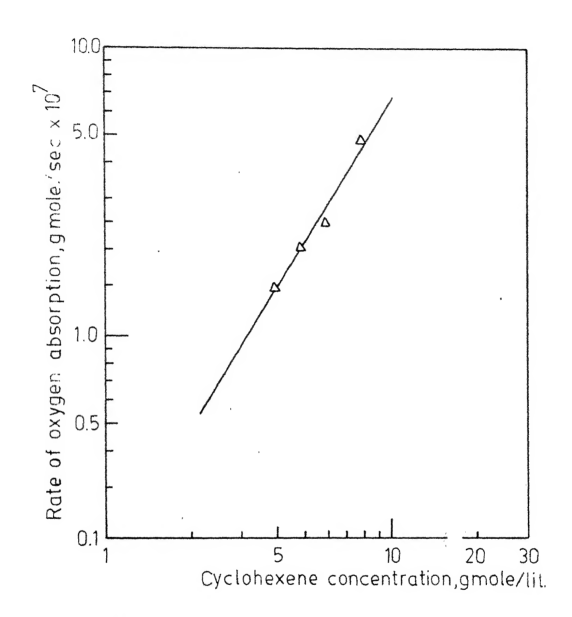


Fig. 8 - Rate dependence with respect to cyclohexene concentration for 0.22 g MnO<sub>2</sub> per 1ml. of cyclohexene at 60°C.

#### Tetralin:

In case of tetralin a catalyst ratio of 0.9 g/ml was employed. This was again chosen from the inhibiting region (Figure 5). The oxygen absorption data are shown in Table C.2 (Appendix C). In Figure 9 rates of oxygen absorption are plotted against hydrocarbon concentrations.on a log-log plot. The order with respect to hydrocarbon concentration was found to be 2.1.

#### 3.4 Decomposition of Hydroperoxide:

The decomposition reactions of the hydroperoxides of cumene, cyclohexene and tetralin were studied for various catalyst ratios at the temperatures of 80, 60 and 65°C, respectively. The decomposition products observed were alcohol, ketone and water. The data are presented in Tables A.3, B.3 and C.3. Figures 10-12 represent the percentage of hydroperoxide decomposed against various catalyst ratios using a single initial hydroperoxide concentration in each case. It is interesting to observe that the percentage decompostion increases with the increasing catalyst ratio upto the break-in-point and then shows a decrease on further increase in the catalyst ratio.

#### 3.5 Reaction Mechanism:

The experimental observations made during the study of oxidation of cumene, cyclohexene and tetralin may be

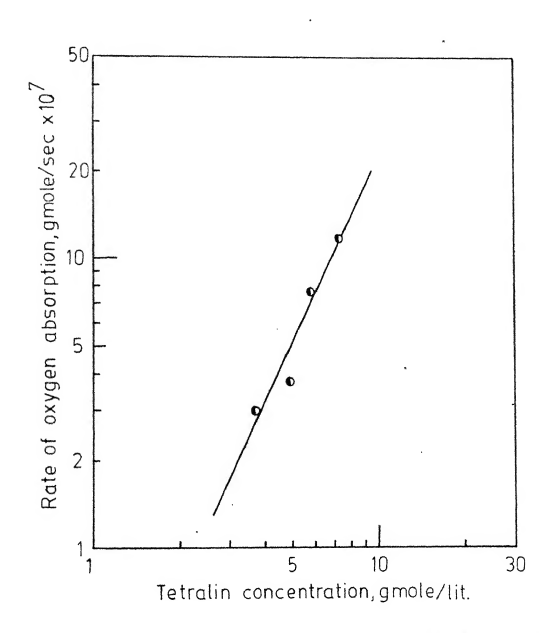
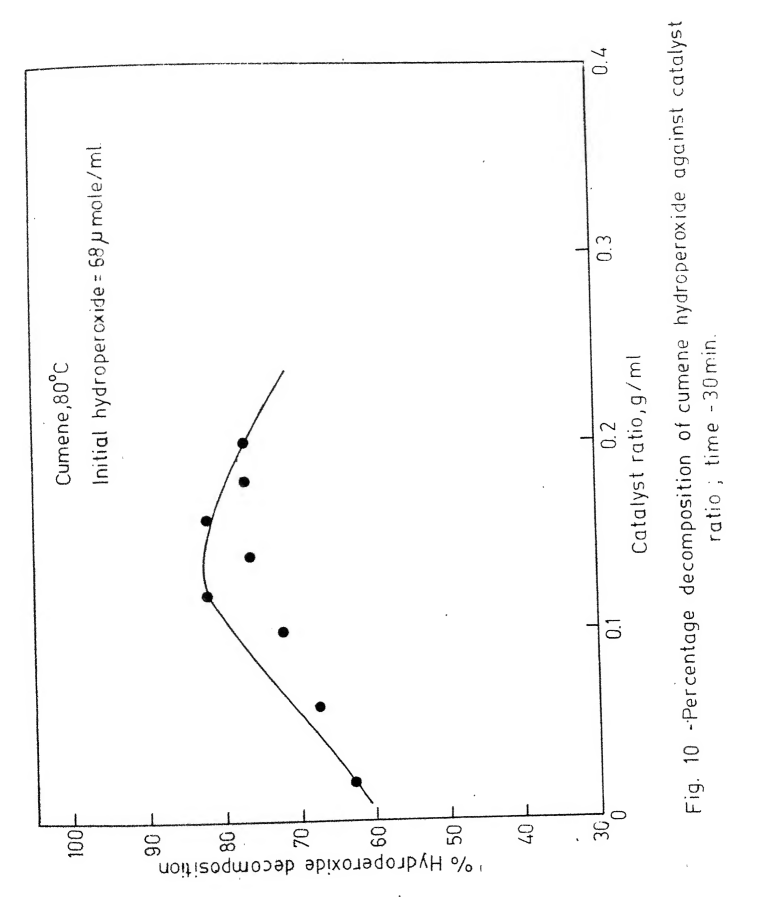
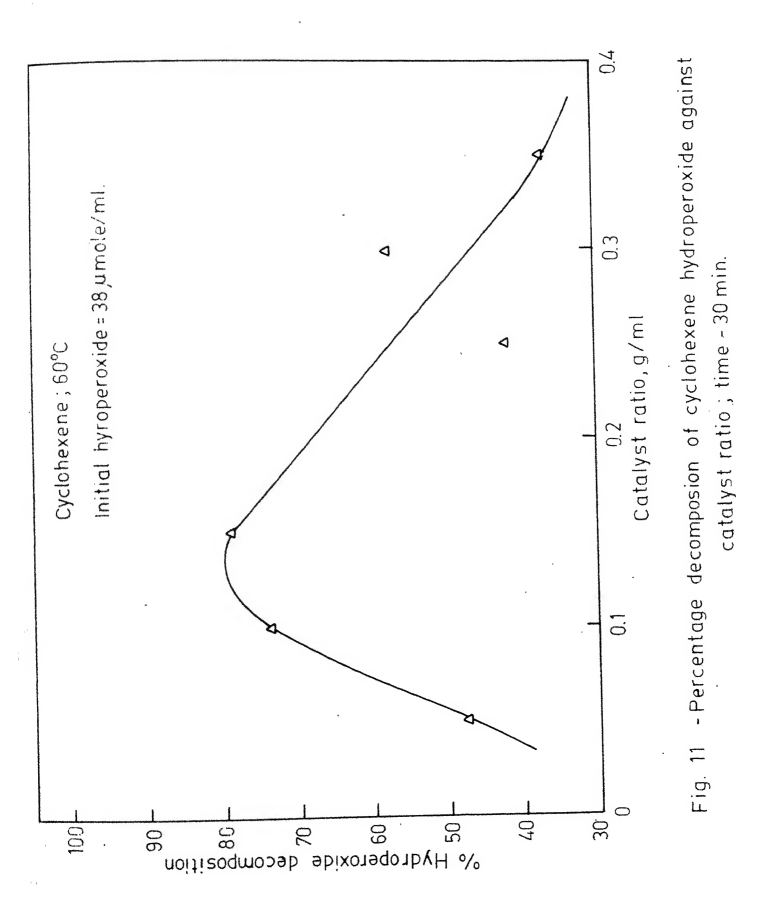
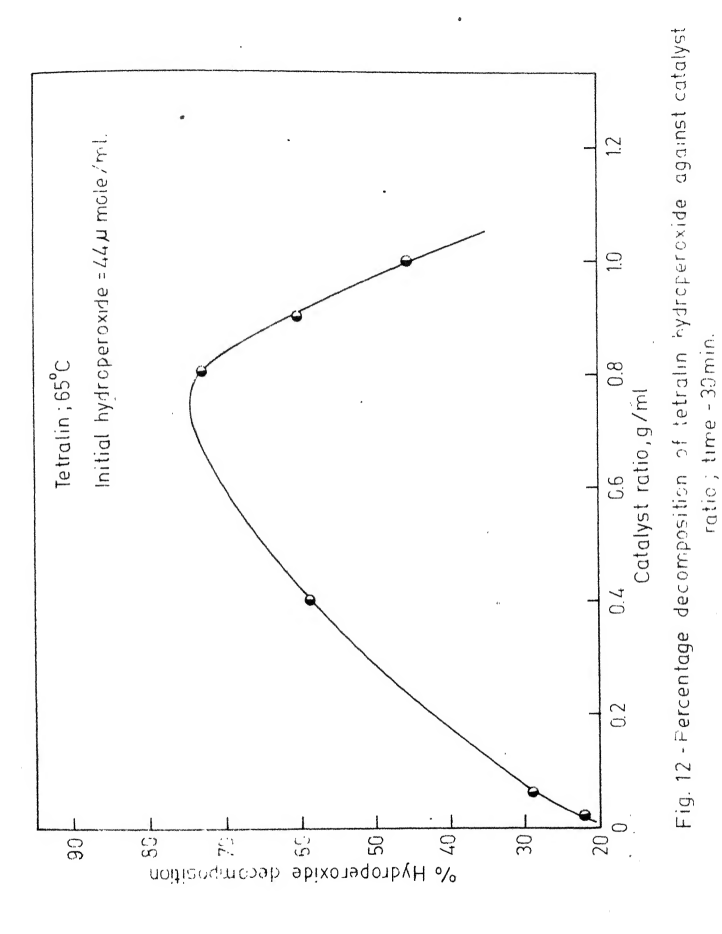


Fig. 9 -Rate dependence with respect to tetralin concentration for 0.9 g MnO<sub>2</sub> per 1 ml. of tetralin at 65°C.







summarized in general as follows:

- (a) For lower catalyst ratios the rate of oxidation increases with increasing catalyst ratio. higher catalyst ratios, there is a decline in the rate of oxidation which is referred to as the inhibiting region.
- (b) For the catalyst ratios in the inhibiting region, the rate of oxidation is approximately second order with respect to the hydrocarbon concentration.
- (c) The decomposition of hydroperoxide in all the three cases increases with increasing catalyst ratio, passes through a maximum and then declines suddenly.

Based on the observations made in the present study and from the established facts that, before the limiting region, the rate dependence is 0.5 order with respect to catalyst weight and first order with respect to hydrocarbon concentration, a general mechanism as proposed by Neuburg et al. [7] is described below:

### Initiation |

ROOH + M 
$$\frac{k_1}{}$$
 (ROOH...M) (1)  
(ROOH...M)  $\frac{k_2}{}$  RO' + M...OH (2)  
ROOH + M...OH  $\frac{k_3}{}$  RO' + H<sub>2</sub>O + M (3)  
RO' + RH  $\frac{k_4}{}$  ROH + R' (4)

$$(ROOH...M) \xrightarrow{k_2} RO^{\bullet} + M...OH^{-}$$
 (2)

ROOH + M...OH 
$$\longrightarrow$$
 RO<sub>2</sub> + H<sub>2</sub>O + M (3)

$$RO^{\bullet} + RH \xrightarrow{\Lambda^{4}} ROH + R^{\bullet} \tag{4}$$

Propagation:

$$R^{\bullet} + O_{2} \xrightarrow{k_{5}} RO_{2}^{\bullet}$$

$$RO_{2}^{\bullet} + RH \xrightarrow{k_{6}} ROOH + R^{\bullet}$$
(5)

$$RO_2^{\bullet} + RH \xrightarrow{RG} ROOH + R^{\bullet}$$
 (6)

Termination:

$$RO_{2}^{\bullet} + RO_{2}^{\bullet} \xrightarrow{k_{7}} \text{ inactive products} + O_{2} \qquad (7)$$

$$RO_{2}^{\bullet} + M \xrightarrow{k_{9}} \qquad (RO_{2}...M) \qquad (8)$$

$$RO_{2}^{\bullet} + (RO_{2}...M) \xrightarrow{k_{9}} \text{ inactive products} \qquad (9)$$

$$RO_2^{\bullet} + M \qquad \frac{18}{2} \qquad (RO_2 \dots M) \qquad (8)$$

$$RO_2^{\bullet} + (RO_2...M) \xrightarrow{hg}$$
 inactive products (9)

### 3.6 Derivation of Kinetic Expression:

At steady-state, the rate of initiation of free radicals is equal to the rate of consumption of free radicals.

Therefore: Rate of initiation of free radicals

$$R_i = 2 k_7 [RO_2^*]^2 + k_9 [RO_2^*][RO_2^*...M]$$
 (10)

Reaction (8) is under equilibrium.

Therefore: 
$$k_8 = \frac{\begin{bmatrix} ROO \dots M \end{bmatrix}}{\begin{bmatrix} ROO \end{bmatrix} \begin{bmatrix} M \end{bmatrix}}$$
 (11)

$$[ROO...M] = k_8 [RO_2^{\bullet}] [M]$$

$$R_{i} = 2 k_{7} [RO_{2}^{\bullet}]^{2} + k_{8}k_{9} [RO_{2}^{\bullet}]^{2} [M]$$

$$= [RO_2^*]^2 (2 k_7 + k_8 k_9 [M])$$

$$[RO_{2}^{\bullet}] = \left(\frac{R_{1}}{2k_{7} + k_{8}k_{9}[M]}\right)^{\frac{1}{2}}$$
 (12)

Rate of oxygen consumption is given by

$$-\frac{d[0_2]}{dt} = k_6 [R0_2^{\bullet}] [RH] + R_1 - k_7 [R0_2^{\bullet}]^2$$
 (13)

$$= k_{6} \left( \frac{R_{i}}{2 k_{7} + k_{8} k_{9} [M]} \right)^{\frac{1}{2}} [RH] + R_{i} \left( 1 - \frac{k_{7}}{2 k_{7} + k_{8} k_{9} [RH]} \right)$$
 (14)

Assuming formation of oxygen through termination is negligible

$$-\frac{d[O_2]}{dt} = k_6 \left(\frac{R_i}{2 k_7 + k_8 k_9 [M]}\right)^{\frac{1}{2}} [RH]$$
 (15)

If the termination of peroxy radicals on the catalyst surface is negligible compared to biradical termination

$$-\frac{d[O_2]}{dt} = k_6 \left(\frac{R_1}{2 k_7}\right)^{\frac{1}{2}} [RH]$$
 (16)

It may be assumed that  $R_i$  is proportional to the amount of catalyst. This rate expression (Eq. 16) clearly accounts for the observations of 0.5 and 1 order with respect to the catalyst ratio and the hydrocarbon concentration, respectively.

When the limiting rate condition is reached, the rate of formation of hydroperoxide by reaction (6) becomes equal to the rate of decomposition on the catalyst surface,  $R_{\rm d}$ . Hence the net rate of formation of hydroperoxide is

$$\frac{d[ROOH]}{dt} = k_6 \left(\frac{R_i}{2 k_7 + k_8 k_9 [M]}\right)^{\frac{1}{2}} - R_d = 0$$
 (17)

Let 
$$R_i = \alpha R_d$$
 (18)

where,  $\alpha$  is the fraction of hydroperoxide decomposed.

From equations(17) and (18), solving for  $R_i$  give

$$R_{i} = \frac{k_{6}^{2} \alpha^{2} [RH]^{2}}{2k_{7} + k_{8}k_{9}[M]}$$
 (19)

By substituting the value of  $R_i$  in Equation (14), the limiting rate of oxygen consumption becomes

$$-(\frac{d[0_2]}{dt})_{\infty} = \frac{\alpha k_6^2 [RH]^2}{2k_7 + k_8 k_9 [M]} [1 + \alpha(1 - \frac{k_7}{2k_7 + k_8 k_9 [M]})]$$
(20)

For small value of  $\boldsymbol{\alpha}$  the limiting rate of oxidation is given by

$$-(\frac{d[O_2]}{dt})_{co} = \frac{\alpha k_6^2 [RH]^2}{2k_7 + k_8 k_9 [M]}$$
 (21)

Therefore, under limiting condition the oxidation rate must show a second order dependence on the hydrocarbon concentration. The experimental observations for all the three systems cumene, cyclohexene and tetralin give a rate dependence of approximately 2. The observations are, therefore, in good conformity with the rate expression derived. The similarity in the kinetic behaviour of the three systems may be due to formation of resonance stabilized radical species. Therefore, it is envisaged that a single mechanism satisfactorily explains the kinetic behaviour observed for all these systems.

### 4. STATISTICAL ANALYSIS OF THE DATA

In order to further substantiate the model a statistical analysis was done. The statistical treatment involved estimation of parameters for the three systems under study, establishing confidence intervals [14] for the parameters and then a residual analysis [15].

# 4.1 Estimation of Parameters and Their Confidence Intervals:

The parameter estimation was essentially based on the minimization of the sum of squares between the expected and the experimental responses with a suitable convergence criterion. Rosenbrock [16] optimization program was used for the minimization. IBM 1800 and IBM 7044 computers were used. The method is of direct-search type. An initial estimate was used to start with. The program was tested for convergence using Banana function,

$$F = 25 (x_2-x_1)^2 + (1-x_1)^2$$

For a fast convergence and simplification the non-linear model was reduced to a three parameter one as follows:

$$\frac{d[O_2]}{dt} = \frac{b_1 [RH]^2}{b_2 + b_3 [M]}$$

where, 
$$b_1 = \alpha k_6$$
  
 $b_2 = 2 k_7^2$   
 $b_3 = k_8 k_9$ 

The problem was reduced to estimating the three parameters b, b, and b, in each of the three systems.

The parameters thus obtained and their 95 per cent confidence intervals for the three systems are tabulated below. The program listings and the calculation of confidence intervals are included in Appendix D<sub>1</sub>.

System	Parameter Values	95 per cent confidence intervals
Cumene	$b_1 = 2.2588 \times 10^2$	$1.9598 \times 10^{2} \le \beta_{1} \le 2.5578 \times 10^{2}$
	$b_2 = 1.3744 \times 10^{10}$	$1.1924 \times 10^{10} \beta_{2} < 1.5564 \times 10^{10}$
	$b_3 = 1.2048 \times 10^3$	$-1.0600 \times 10^{10} = \beta_{3} < 1.0600 \times 10^{10}$
Cyclohexene	$b_1 = 2.1638 \times 10^2$	$1.7758 \times 10^2 \le \beta_1 - 2.5518 \times 10^2$
	$b_2 = 1.3744 \times 10^{10}$	$1.1274 \times 10^{10} \beta_2 < 2.6214 \times 10^{10}$
	$b_3 = 1.0500 \times 10^3$	$-1.1280 \times 10^{10} \lesssim \beta_3 < 1.1200 \times 10^{10}$
Tetralin	$b_1 = 1.9584 \times 10^2$	$-2.1042 \times 10^3 \le \beta_1 < 2.5058 \times 10^3$
	$b_2 = 3.6280 \times 10^9$	$3.0140 \times 10^{9} \times \beta_{2} \times 4.2419 \times 10^{9}$
	$b_3 = 1.8140 \times 10^9$	$1.1326 \times 10^9 \approx \beta_3 < 2.4954 \times 10^9$

An examination of the confidence intervals indicates that the parameter values are all positive in the confidence region. This is one of the important criteria for model selection.

## 4.2 Residual Analysis:

The residual analysis is a simple but powerful tool to test the adequacy of a model. A residual is the difference

between an observed value and the corresponding predicted value. Using the estimated parameters, rates were predicted (Appendix D3). The residuals were then obtained by subtracting the predicted rates from the observed ones. Figures 13, 14 and 15 show the residual plots against the independent variable viz. hydrocarbon concentration for the systems cumene, cyclohexene and tetralin, respectively. The plots show a random behaviour of the residuals which further substantiates the model.

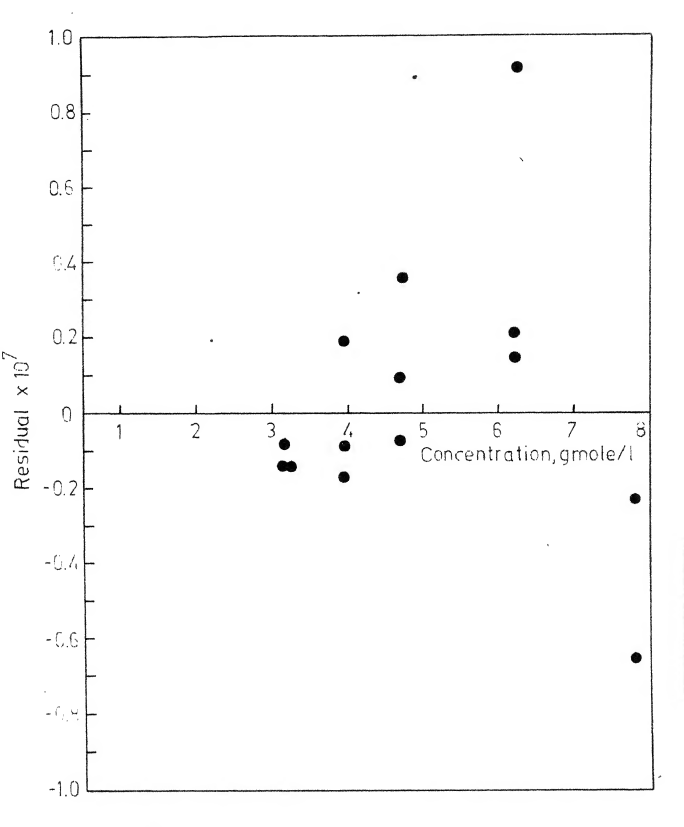


Fig. 13 -Plot of the residuals against the hydrocarbon concentration for the system cumene.



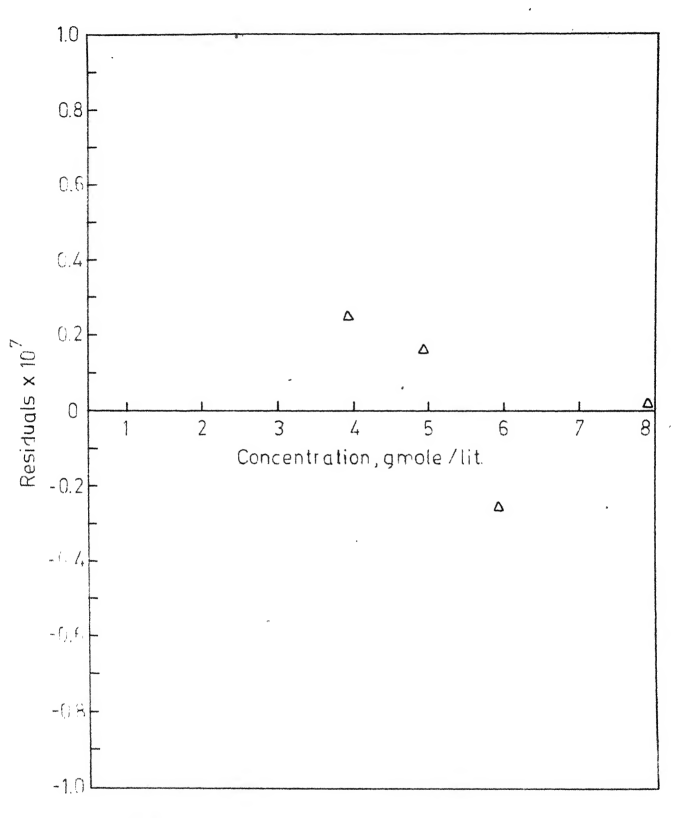


Fig. 14 - Plot of the residuals against hydrocarbon concentration for the system cyclohexene.

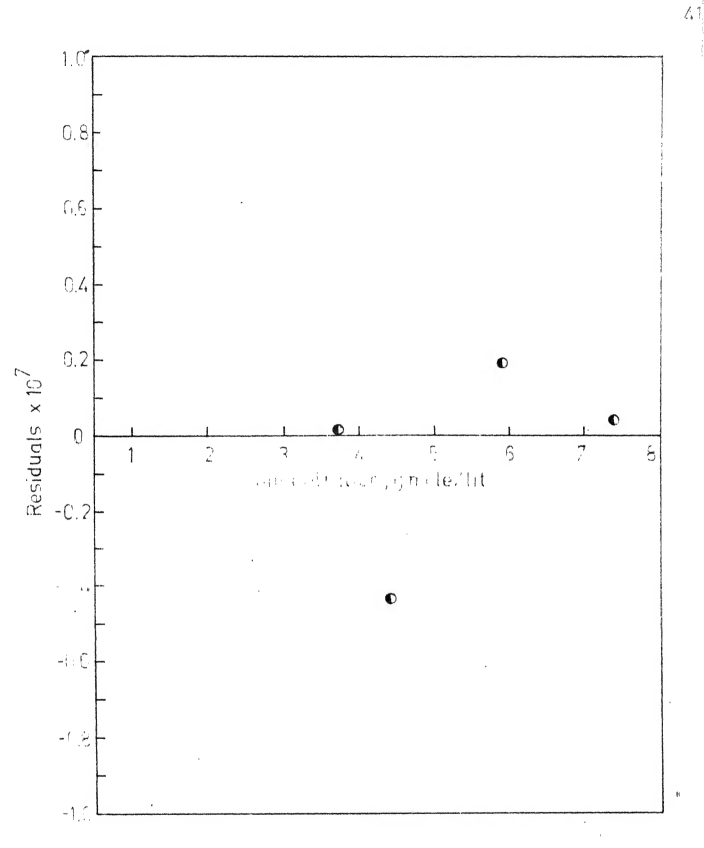


Fig. 15 - Plot of residuals against the hydrocarbon concentro tor the system tetralin.

### 5. CONCLUSIONS

The kinetics of oxidation of the three systems — cumene, cyclohexene and tetralin showed a similar behaviour. In all the three cases it was established that as the catalyst to hydrocarbon ratio was increased, the rate of oxygen absorption increased and after a certain ratio the rate declined suddenly. While an order of one with respect to hydrocarbon concentration was established earlier in the region before the break-in-point, a second order rate dependence on hydrocarbon concentration was obtained in the inhibiting region. The second order dependence in the inhibiting region supports the degenerate chain-branching mechanism.

The observed data were given a statistical treatment to test the adequacy of the model. The parameters were estimated using Rosenbrock's optimization method. The confidence intervals were also estimated. The parameters were all positive in the confidence region. Finally, a residual analysis showed a random nature with respect to the independent variable, hydrocarbon concentration. The positive nature of the parameters and the random behaviour of the residuals substantiate the model derived on the basis of degenerate chain-branching mechanism. Thus, such model, in general, accounts for the reactions where hydroperoxide forms and propagates the chain reaction.

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APPENDIX A
CUMENE RUNS

TABLE A-1: OXYGEN ABSORPTION VERSUS TIME AT DIFFERENT Mno  $_{\rm 2}$  CATALYST WEIGHTS AT 80  $^{\rm o}{\rm C}$ 

Run No.	Catalyst ratio, g/ml	Time, min.	Oxygen consumption (g mole/ml)x10 <sup>4</sup>	Initial rate g mole/sec x 10 <sup>7</sup>
C-1	0.02	0	0	1.934
		5	0.602	
		10	1.160	
		15	1.722	
		20	2.250	
		25	2.940	
		30	3.364	
		35	3.785	
		40	4.080	
C-2	0.06	0	0.000	3.600
		5	1.080	
		10	2.065	
		15	2.940	
		20	3.505	
		25	3.880	
•		30	4.020	
		35	4.235	
		40	4.315	

Table A-1 (contd)

Run No.	Catalyst ratio, g/ml	Time, min.	Oxygen consumption, (g mole/ml)x 10 <sup>4</sup>	Initial rate g mole x107 sec
C-3	0.10	0	0.000	4.170
		5	1.340	
		10	2.412	
		15	3.205	
		20	3.719	
		25	4.100	
		30	4.290	
		35	4.480	
C-4	0.14	0	0.000	5.170
		5	1.763	
		10	3.055	
		15	3.840	
		20	4.310	
		25	4.520	
		30	4.660	
C <b>-</b> 5	0.160	0	0.000	4.800
		5	1.384	
		10	2.550	
		16.5	3.449	
		20	3.719	
		25	4.025	
		. 30	4.220	
		35	4.340	
		40	4.440	

Table A-1 (contd)

Run No.	Catalyst ratio, g/ml	Time, min.	Oxygen consumption, (g mole/ml)x10 <sup>4</sup>	Initial rate, g mole x 107 sec
C-6	0.18	0	0.000	4.380
		5	1.276	
		10	2.389	
		15	3.110	
		20	3.460	
		25	3.745	
C-7	0.20	0	0.000	2.866
		5	0.814	
		10	1.850	
		15	2.429	
		20	2.879	
		25	3.1.40	

TABLE A-2: OXYO EN ABSORPTION VERSUS TIME FOR DIFFERENT CUMENE CONCENTRATIONS (LIMITING ZONE)

Run No.	Catalyst ratio, g/ml	Cumene concen- tration, g mole/lit	Time, min.	Oxygen absorption, g mole x 104	Initialrate, g mole x 107 sec
C-8	0.16	7.82	0	0.000	4.800
			5	1.384	
			10	2.550	
			16.5	3.449	
			20	3.719	
			25	4.025	
			30	4.220	
			35	4.340	
			40	4.410	
C-9		6.26	0	0.000	4.130
			5	1.078	1
			10.5	2.015	
			17	2,905	
			20	3.175	
			25	3.519	
			30	3.685	
			35	3.920	
C-10	)	4.69	0	0.000	1.731
			5	0.414	
			10	0.909	
			15	1.528	

Table A-2 (contd)

Run No.	Catalyst ratio, g/ml	Cumene con- centration, g mole/lit	Time, min.	Oxygen absor- ption, g mole x 10 <sup>4</sup>	Initial rate, g mole x10
			20	1.928	
			25	2.514	
			30	2.930	
			35	3.392	
			40	3.760	
C-ll		3.99	0	0.000	1.142
			5	0.299	
			10	0.644	
•			15	1.018	
			20	1.390	
			25	1.738	
			30	2.049	
			35	2.375	
			4 Q	2.680	
C-12	0.16	3.13	0	0.000	0.667
			5	0.197	
			10	0.380	
			15	0.583	
			20	0.814	
			25	1.031	
			30	1.261	
			35	1.471	
			40	1.710	

Table A-2 (contd)

Run No.	Catalyst ratio, g/ml	Cumene con- centration, g mole/bit	Time, min.	Oxygen absor- ption, g molex 104 ml	Initial rate, g molex 107
C-13	0.18	7.82	0	0.000	4.380
			5	1.276	
			10	2.389	
			15	3.110	
			20	3.460	
*			25	3.745	
C-14		6.26	0	0.000	3.430
			5	1.004	
,			10	1.940	
			15	2.640	
			20	3.230	
			25	3.550	
			30	3.810	
			35	Z.990	
			40	4.120	
C-15		4.69	0	0.000	2.166
			5	0.634	
			10	1.229	
			15	1.754	i i
			20	2.209	
			25	2.691	
			30	and the contract of the contra	
			35	3.520	IBRARY
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Table A-2 (contd)

Run No.	Catalyst ratio, g/ml	Cumene con- centration, g mole/lit	Time, min.	Oxygen ab- sorption, g molex104	Initial rate, g mole x10
C-16	0.18	3.99	0	0.000	1.220
			5	0.345	
			10	0.732	
	-		15	1.173	
	,		20	1.532	
			25	1.864	
			30	2.154	
			35	2.470	
			40	2.749	
C-17		3.13	0	0.000	0.667
			5	0.207	
			10	0.394	
			15	0.593	
			20	0.828	
			25	1.008	
			30	1.256	
			35	1.470	
			40	1.711	
C-18	0.20	6.26	0	0.000	3.365
			5	0.966	
			10	2.070	
			15	2.925	
			20	3.570	

Table A-2 (contd)

Run No.	Catalyst ratio, g/ml	Cumene con- centration, g mole/lit	Time, min.	Oxygen ab- sorption, g mole x10 <sup>4</sup>	Initial rate, g mole <sub>x</sub> 10 <sup>7</sup>
			25	3.990	
			30	4.265	
			35	4.400	
			40	4.500	
C-19	0.20	4.69	0	0.000	1.900
			5	0.566	
			10	1.160	
			20	2.235	
			25	2.745	•
			30	3.130	
			35	3.465	
			40	3.625	
C-20		3.99	0	0.000	1.500
	•		5	0.469	
			10	0.897	
			15	1.241	
			20	1.531	
			25	1.884	
			30	2.090	
			35	2.499	
			40	2.840	

Table A-2 (contd)

Run No.	Catalyst ratio, g/ml	Cumene con- centration, g mole/lit	Time, min.	Oxygen ab- sorption, g mole x104	Initial rate, g mole <sub>x10</sub> 7
C-21		3.13	0	0.000	0.716
			5	0.176	
			10	0.401	
			15	0.704	
			20	0.870	
			25	1.090	
			30	1.318	
			35	1.657	
			40	1.815	
Annual Section Section 40000 and				_	

TABLE A-7: CUMENE HYDROPEROXI)E DECOMPOSITION AT 80°C FOR DIFFERENT CATALYST (MnO2) RATIOS

Run No.	Catalyst ratio,	Initi	al Concel	ntı	Final	Final Concentration,	ation,	Hyderoperoxide
	g/m	HPR <sup>+</sup>	KET <sup>++</sup>	$A \operatorname{LC}^{lpha}$	HPR	KET	A_LC	110 Hz 04 m000 s
G-22	0.02	68.1	06	73.4	25.1	70.0	77.8	63.1
G-23	90.0	68.1	90	73.4	22.1	0.06	78.5	67.5
G-24	0.10	68.1	90	73.4	18.0	0.06	9.57	72.4
0-25	0.12	68.1	90	73.4	12.3	190.0	78.9	82.0
0-26	0.14	68.1	96	73.4	16.0	150.0	80.2	76.5
G-27	0.16	68.1	90	73.4	12.3	130.0	79.3	82.0
G-28	0.18	68.1	90	73.4	22.3	100.0	78.4	67.3
G-29	0.20	68.1	90	73.4	22.3	100.0	78.4	67.3
+Hydro	+ Hydroperoxide	++Ketone		αAlcohol			e Cristian inglanding springs, makes, const. a.o.o.	

APPEMDIX B

CYCLOHEXENE RUNS

TABLE B-1: OXYGEN CONSUMPTION VERSUS TIME FOR DIFFERENT MnO CATALYST WEIGHTS AT  $60^{\circ}\text{C}$ 

Run No.	Catalyst ratio, g/ml	Time, min.	Oxygen absor- ption, g mole ml	Initial rate, g mole x10 <sup>7</sup> sec
C-30	0.08	0	0.000	1.335
		5	0.103	
		10	0.937	
		15	1.756	
		20	2.270	
C-31	0.12	0	0.000	2.000
		5	0.571	
		8	0.780	
		10	0.908	
		15	1.274	
C-32	0.18	0	0.000	2.500
		5	0.586	
		10	0.835	
		15	1.180	
		20	1.370	
C-33	0.20	0	0.000	3.000
		5	0.880	
		10	1.610	
		15	1.990	
		20	2.230	

Table B-1 (contd)

Run No.	Catalyst ratio, g/ml	Time, min.	Oxygen absor- ption, g mole x 10 <sup>4</sup>	Initial rate, g mole x 10 <sup>7</sup> sec
C-34	0.32	0	0.000	2.400
		5	0.464	
		10	1.215	
		15	1.760	
		20	2.125	
C-35	0.40	0	0.000	1.500
		5	0.461	
		10	1.025	
		15	1.464	
		20	1.859	

TABLE B-2: OXYGEN CONSUMPTION VERSUS TIME FOR DIFFERENT CYCLO-HEXENE CONCENTRATIONS (LIMITING ZONE) AT 60°C

Run No.	Catalyst ratio, g/ml	Cyclohexene concentration, g mole/lit	Time, min.	Oxygen consumption g mole x104	Initial rate g mole x10 <sup>7</sup> sec
C-36	0.22	7.880	0	0.000	4.90
			5	1.377	
			10	2.254	
			15	2.750	
			20	3.149	
			25	3.380	
			30	3.540	
C-37	5.92	5.920	0	0.000	2.50
			5	0.615	
			10	1.464	
			15	2.250	
			20	2.855	
C-38		4.930	0	0.000	2.08
			5	0.659	
			10	1.518	
			15	1.918	
	•		20	. 2.430	
			25	2.955	
			30	3.395	

Table B-2 (contd)

Run Mo.	Catalyst ratio, g/ml	Cyclohexene concentration, g mole/lit	Time, min.	Oxygen con- sumption, g molex104	Initial rate g mole x107 sec
C-39		3.945	0	0.000	1.50
			5	0.439	
			10	0.922	
			15	1.428	
			20	1.889	
			25	2.320	
			30	2.755	

TABLE B-3: CHYCLOHEXENE HYDROPEROXIDE DECOMPOSITION AT 60°C FOR DIFFERSNT CATALYST (MnO<sub>2</sub>) RATIOS

		TORREST RETURNATION AND CONTRACT CONTRA	Andreas of the Constitution Committee of the Constitution.	THE REPORT OF THE PARTY OF THE PARTY.	S. March Constituted and Acres of March States	STATE AND THE PARTY AND TOTAL OF THE PARTY O	AND THE PERSON NAMED IN COLUMN TWO IS NOT THE OWNER.	MARKAMAN MENUNCATA PARAMAN MARKAMAN MENUNCAK MENANGKA PETANG ARAB ARAB PENANGKAN PETANGKAN PERANGKAN PERANGKAN
Run No.	Catalyst ratio	Initial C	Concentration,	ation,	Final	Final Concentration	ation	Per cent Hydropero-
	g/ml	HPR	KET	ALC	HPR	KLT	ALC	xide decomposition
0-40	0.05	38	1490	ا	20	1180	Q.	4.7.4
0-41	0.10	38	1490	ا	10	1020	τΩ	73.6
0-42	0.15	38	1490	ا	ώ	550	ا	0.67
C-43	0.25	38	1490	Q I	22	1100	م	42.0
C-44	0.30	38	1490	ا	16	006	وم	0 7.7
0-45	0.35	38	1490	رم ا	24	1360	٦ ع	36.9
Part of the Part of Spirits of the Part of	COMPTO T THE MELTINESTEEN THE MEST THE THE THE THE THE THE THE THE THE TH	A THE STATE OF THE PROPERTY OF THE PERSON	TO A SECURE DEPENDENT MANAGEMENT OF THE PROPERTY OF THE PROPER	BOY IL THE PROPERTY METERS THE	en ann it – 16 filestandringsjeren en ann			\

 $^{\mathrm{b}}$  Not estimated

APPENDIX C

TETRALIN RUNS

TABLE C-1: OXYGEN CONSUMPTION VERSUS TIME FOR DIFFERENT MnO<sub>2</sub>

Run No.	Catalyst ratio, g/ml	Time, min.	Oxygen consumption, g mole x 10 <sup>4</sup>	Initial rate g mole x 107 sec
C-47	0.2	0 5 10 15 20	0.000 1.081 1.750 2.300 2.800	3.6
C-48	0.4	0 5 10 15 20	0.000 2.065 3.500 4.500 5.100	6.6
C-49	0.8	0 5 10 15 20	0.000 1.514 2.600 3.300 4.000	10.1
C-50	0.9	0 5 10 15 20	0.000 2.880 4.110 5.190 5.980	9.6
C-51	1.0	0 5 10 16 20	0.000 0.180 0.450 1.081 1.587	0.600

Table C-2 (contd)

Run No.	Catalyst ratio, g/ml.	Tetralin con- centration, g mole/lit	Time, min.	Oxygen con- sumption, g molex 104 ml	Initial Rat g mole x 1( sec
	- Commence of the Commence of		15	2.129	
			20	2.705	
			25	3.210	
			30	3.675	
			35	4.110	
			40	4.470	
C-55		3.69	0	0.000	3.00
			5	0.829	1
			10	1.298	1
			15	1.730	
	,		20	2.125	
			25	2.625	
			30	2.685	1 1
			35	3.245	t t
			40	3.605	

TABLE C-3: TETRALIN HYDROPEROXIDE DECOMPOSITION AT 65°C FOR DIFFERENT CAPALYST (MnO<sub>2</sub>) RATIOS

Run No.	Catalyst ratio, g/ml	Initial µ HPR	Initial Concentration  µ mole/ml  HPR  KET  ALC	ation	Final ( (30 m HPR	Final Concentration (30 min run) pmole HPR KET ALC	Concentration min run) pmole/ml KET	Per cent Hydroperoxid decomposition
77.7	0.02	44.2	ರ	ا م	34.5	ದ	q Q	22.0
	90.0	44.2	ದೆ	ا	12.7	ದ	ا <sup>ث</sup>	28.8
, , , , , , , , , , , , , , , , , , ,	0,40	44.2	ರೆ 	۱	18.1	ದೆ [	ا <sup>م</sup>	59.1
C ()	08,0	44.2	<sup>ಹ</sup> !	ا	12.0	ದ	۱	72.9
α-60	0.90	44.2	g I	۱	17.6	ර I	٦٩	60.2
C-61	1,00	44.2	್ (	ا	24.1	g !	ر م	45.5
	A SA PRESENTATION OF SERVICE SERVICES SERVICES.	PETERS OF THE PE	SERVICE PARTY PART	The best and the second of the	-derothen, desten krapting å n d de fless den de side.	Men semante e manufacture semantes.	Labeladah, o' Add. Teoriti mila, N. S.	A STATE OF THE PARTY OF THE PAR

 $^{
m a}_{
m Detected}$  in infrared spectra

<sup>&</sup>lt;sup>b</sup>Not estimated

APPENDIX D

STATISTIC L ANALYSIS

ROS00390

ROS00400

```
69
     II = II + 1
     D \times (II) = (X(1) * RH(II) * * 2) / (2.* X(2) + X(3) * CAT(I))
      D02CJ=1, NN
21
     CUNTINUE
     FUNXCN=0.0
     DD30 I=1.N
     FUNXCH=FUNXON+(10.**7*DX(I)-DX1(I))**2
30
     CONTINUE
     RETURN
     END
С
                                                                ROSOCCIO
C
 C
                                                                R0S00030
  EVALUATION OF MAXIMUM/MINIMUM OF A FUNCTION BY ROSENBROCK METHOD
С
                                                                RUS00040
  V* COMPONENTS OF ORTHOGONAL VECTORS
С
                                                                ROS00050
C
  SUBPREGRAM FUNXON DEFINES FUNCTION F(X,Z) AS F(0,Z)
                                                                ROS00060
C
  X* UNKNOWN VARIABLES
                                                                R0S00070
C
  Z* KNOWN PARAMETERS
                                                                R0S00080
C
  N* NUMBER OF UNKNOWN VARIABLES,X
                                                                R0S00090
C
  NN* NUMBER OF KNOWN PARAMETERS, Z
                                                                ROS00100
C
  NS* NUMBER OF STAGES AFTER WHICH CALCULATION SEQUENCE IS TERMINATED
                                                                ROS00110
С
  MAXMIN = ZERO FOR MAXIMIZATION AND ONE FOR MINIMIZATION
                                                                R0500120
C
  LIMIT = ZERO FOR FINITE LIMIT AND ONE FOR NO FINITE LIMIT
                                                                ROS00130
C
  ALPHA* A POSITIVE MULTIPLYING FACTOR FOR INCREASING A SUCCESSFUL STEPROSO0140
C
  BETA* A POSITIVE MULTIPLYING FACTOR FOR DECREASING UNSUCCESSFUL STEPSROSOO 150
C
  E* INTIAL STEP SIZE AT THE BEGINNING OF EACH STAGE
                                                                RDS00160
U
  FL* VALUE OF FINITE LIMIT
                                                                ROS00180
     FORMAT(515,4F10.6)
1
                                                                RUS00190
     READ 2, (X(I), I=1, N), ((V(I,J), J=1, N), I=1, N)
                                                                ROS00200
2
      FORMAT(8F10.6)
                                                                R0S00210
     PRINT 3, N, NS, E, ALPHA, BETA
                                                                R0S00220
     3
                        NUMBER OF VARIABLES (N) *, 16, 12X, 1H*/1X, 3ROSQ0240
    1,1H*,48X,1H*/1X,31H*
           NUMBER OF STAGES (NS)
    21H*
                                  *, I6, 12X, 1H*/1X, 31H*
                                                       INITIAL TRROSO0250
    3IAL STEP (E)
                  *,F10.6,8X,1H*/1X,12H*
                                         ALPHA *, F10.6, 3X, 6HBETA *ROSO0260
    5**********
                                                                ROS00280
     IF(MAXMIN.EQ.0) GO 10 11
                                                                R0500290
     IF(LIMIT.EQ.O) GO TO 13
                                                                R0S00300
     PRINT 10
                                                                ROS00310
     GO TO 17
                                                                ROS 00320
4
     CALL MINMAX(N,NS,MAXMIN,LIMIT,ALPHA,BETA,E,FL,X,V,Z)
                                                                R0S00330
     GO TO 8
                                                                ROS00340
5
     READ 6, (Z(I), I=1,NN)
                                                                ROS 00350
     FORMAT(8F10.6)
                                                                ROS00360
6
     PRINT 7,(Z(I),I=1,NN)
                                                                ROS00370
7
     FORMAT(//7X,3HZ *,8F15.8)
                                                                RDS 00380
```

GO TO 4

PRINT 9

8

```
9
      FORMAT(//47x,35HCOMPLETED REQUIRED NUMBER OF STAGES//)
                                                                              RDS00410
      STOP
                                                                              ROS00420
10
      FORMAT(1HO, 33HMINIMIZATION WITH NO FINITE LIMIT)
                                                                              RUS00430
11
      IF(LIMIT.EQ.O) GO TO 15
                                                                              ROS 00440
      PRINT 12
                                                                              ROS 00450
      GU TO 17
                                                                              ROS00460
12
      FORMAT(1HO, 33HMAXIMIZATION WITH NO FINITE LIMIT)
                                                                              RUS00470
13
      PRINT 14, FL
                                                                              RDS 00480
      GO TO 17
                                                                              ROS 00490
14
      FORMAT(1HC, 35HMINIMIZATION WITH FINITE LIMIT FL =, F1C.6)
                                                                              RUS00500
15
                                                                              ROS00510
      PRINT 16, FL
16
      FORMAT(1HO,35HMAXIMIZATION WITH FINITE LIMIT FL =,F?G.6)
                                                                              RDS 005 20
17
      IF (NN-GT-0) GO TO 5
                                                                              ROS 00530
                                                                               R0S00540
      GO TO 4
      END
                                                                              ROS00550
C
                                                                              ROS 00 5 60
                                                                              ROS 0 U 5 70
      SUBROUTINE MINMAX(N, NS, MAXMIN, LIMIT, ALPHA, BETA, E, FL, X, V, Z)
      DIMENSION A(40,40),B(40,40),C(40,40),P(40,40),V(40,40)
                                                                              ROS 00 5 80
      DIMENSION D(40), D(40), R(40), X(40), Y(40), Z(40)
                                                                              ROS00590
                                                                               RDS00600
      S = E
                                                                               ROSO061U
      L = 1
                                                                               ROS00620
      NSTAGE=-1
      NTRIAL=-1
                                                                               ROS00630
      NSUXES=0
                                                                               RDS 00640
                                                                               ROS00650
      NFEL UR = 0
                                                                               RUS00660
      00 1 I=1,N
                                                                               ROS00670
      D(I) = 0.0
      O(I) = X(I)
                                                                               RDS00680
                                                                               RDS00690
      R(I) = 0.0
                                                                               ROSO07C0
      Y(I) = 0.0
                                                                               R0500710
      DO 1 J=1,N
                                                                               R0S00720
      A(I,J)=0.0
      B(I,J)=0.0
                                                                               RUS 00730
                                                                               R0S00740
      C(I,J) = 0.0
                                                                               R0S00750
      P(I,J)=0.0
                                                                               ROS00760
2
      F = FUNXON(O,Z,N)
                                                                               RUS00770
      IF(NTRIAL.LT.O) GO FO 9
                                                                               R0S00780
      IF(MAXMIN.EQ.O) GO TO 15
                                                                               R0S00790
      IF(LIMIT.EQ.0)
                        GO TO 18
                                                                               R0S00800
                        GO TO 13
3
       IF (F.LE.FM)
       IF(NSTAGE.LT.O) GO TO 5
                                                                               RDS00810
4
                                                                               R0S00820
      NFELUR=NFELUR+1
       IF(NSUXES.GT.O) GO TO 21
                                                                               R0S00830
                                                                               R0500840
5
      E =-BETA*E
                                                                               RDS00850
      NTRIAL=NTRIAL+1
                                                                               ROS00860
       Y(L) = E
                                                                               R0S00870
      DO 8 I=1,N
                                                                               ROS00880
       O(I) = 0.0
                                                                               R0S00890
       DO 7 J=1,N
                                                                               ROS00900
```

(L) Y \* (L, I) V + (I) D = (I) D

7

ROS01390

ROS01400

```
\cap (I) = O(I) + X(I)
8
                                                                               ROS00910
      SO TO 2
                                                                               ROS00920
9
      NTRIAL=NTRIAL+1
                                                                               ROS 00930
      PRINT 10, F, (X(I), I=1, N)
                                                                               RDS00940
      FORMAT(//1X,21H **STARTING VALUES**//7X,3HF *,F15.8//7X,3HX *,
10
                                                                               RDS 009 50
     18F15.8)
                                                                               ROS00960
      PRINT 11
                                                                               RDS 009 70
      FORMAT(//7X,3HV *)
1
                                                                               ROS00980
      PRINT 12, ((V(I,J),J=1,N),I=1,N)
                                                                               R0S00990
12
      FORMAT(10X,4F15.8/)
                                                                               ROS01000
13
                                                                               ROS01010
      IF (NSTAGE-LT-0) GO TO 19
                                                                               ROS 01020
      D(L) = D(L) + Y(L)
                                                                               ROS 01030
      DU 14 I=1,N
                                                                               ROS 0 1 0 4 0
14
      X(I)=O(I)
                                                                               ROS 01050
      NSUXES=NSUXES+1
                                                                               ROS 01 0 60
      IF(NFFLUR.GT.O) GO TO 21
                                                                               RUS 01070
      E = AL PHA *E
                                                                               ROS 01080
      GO TO 6
                                                                               ROS 01090
15
      IF(LIMIT.EQ.O)
                        GO TO 16
                                                                               ROS01100
      GO TO 17
                                                                               ROS01110
                        GO TO 17
16
      IF(F.LE.FL)
                                                                               ROS01120
      GO TO 4
                                                                               ROS01130
17
      IF(F.GE.FM)
                        GO TO 13
                                                                               RUS01140
      30 TO 4
                                                                               ROS 0 1 1 50
      IF(F.GE.FL)
                        GO TO 3
18
                                                                               RDS 011 60
      GO TO 4
                                                                               ROS01170
19
      NSTAGE=NSTAGE+1
                                                                               ROS01180
      GD TO 6
                                                                               ROS01190
21
      Y(L) = 0.0
                                                                               ROS01200
      L=L+1
                                                                               ROS01210
      IF(L.GT.N)
                        GO TO 23
                                                                               ROS01220
22
      F = S
                                                                               ROS01230
      NSUXES=0
                                                                               RUS01240
      NFELUR=0
                                                                               ROS 01 250
      SO TO 6
                                                                               ROS01260
23
      NSTAGE=NSTAGE+1
                                                                               ROS01270
      CALL VECTOR (A,B,C,P,V,D,R,N)
                                                                               RDS 01 280
      TEMP1=R(1)
                                                                               ROS01290
      TEMP2=0.0
                                                                               ROS01300
      DO 24 I=1,N
                                                                               ROS01310
      TEMP2=TEMP2+A(I,2) **2
24
                                                                               ROS01320
      TEMP2=SQRT(TEMP2)/TEMP1
                                                                               ROS01330
      PRINT 25, NSTAGE, NTRIAL, F, TEMP1, TEMP2, (X(I), I=1, N)
                                                                               ROS01340
25
      FORMAT(//21H AT THE END OF STAGE , 16,9H AND
                                                           , I6,9H
                                                                    TRIALS//7RDS01350
     1X,3HF *,F15.8//2X,8HMAG A1 *,F15.8,5X,14HMAG A2/MAG A1*,F15.8//7X,ROSO1360
     23HX *,8F15.8)
                                                                               ROS 01 370
      PRINT 26
                                                                               ROS01380
```

26

FORMAT(//7X, 3HV \*)

PRINT 27, ((V(I,J),J=1,N),I=1,N)

ROS01900

```
27
      FORMAT(10X,4F15.8/)
                                                                                ROS01411
      IF (NSTAGE . EQ.NS) GO TO 29
                                                                                ROS01421
      no 28 I=1,N
                                                                                RUS0143(
                                                                                ROS0144(
      D(I) = 0.0
      R(I) = 0.0
                                                                                ROS0145(
      Y(I) = 0.0
                                                                                ROS01460
      DO 28 J=1,N
                                                                                ROS01470
      \Lambda(I,J)=G.0
                                                                                ROS01480
      B(I,J)=0.0
                                                                                ROS01490
      C(I,J) = 0.0
                                                                                ROS01500
      P(I,J)=9.0
                                                                                ROS01510
28
                                                                                ROS01520
      L=1
      GO TO 22
                                                                                ROS01530
29
      RETURN
                                                                                ROSG1540
                                                                                ROS01550
      FND
                                                                                ROS01560
C
                                                                                ROS 01570
      SUBROUTINE VECTOR(A,B,C,P,V,D,R,N)
                                                                                ROS01580
      DIMENSION A(40,40),B(40,40),C(40,40),P(40,40),V(40,40)
      DIMENSION D(40),R(40)
                                                                                ROS01590
      DO 1 J=1,N
                                                                                ROS01600
      DO 1 I=1,N
                                                                                ROS01610
                                                                                ROS01620
      00 1 K=J,N
                                                                                ROS01630
1
      A(I,J)=A(I,J)+D(K)*V(I,K)
      DO 2 I=1,N
                                                                                RDS 0 1 6 4 0
                                                                                ROS 01650
      B(I,1)=A(I,1)
      R(1) = R(1) + B(I,1) **2
                                                                                ROS 01660
2
                                                                                ROS01670
          R(1) = SQRT(R(1))
                                                                                ROS 01680
      DO 3 I=1.N
3
      V(I,1)=B(I,1)/R(1)
                                                                                ROS 01690
                                                                                ROS01700
        DO 8 J=2,N
                                                                                ROS01710
        MJ = J - 1
                                                                                ROS 0 1 7 2 0
        DO 4 K=1, MJ
                                                                                ROS 01730
        DO 4 I=1,N
                                                                                RUS01740
  4
        C(J,K)=C(J,K)+A(I,J)*V(I,K)
                                                                                ROS01750
        DO 6 L=1,MJ
                                                                                RDS 01760
        DO 6 I=1,N
                                                                                ROS 01 7 70
        P(I,J) = P(I,J) + C(J,L) * V(I,L)
                                                                                ROS01780
  6
        B(I,J) = A(I,J) - P(I,J)
                                                                                ROS01790
        DO 7 I=1,N
                                                                                ROS01800
        R(J)=R(J)+B(I,J)**2
  7
                                                                                ROS01810
          R(J) = SQRT(R(J))
                                                                                ROS01820
        DO 8 I=1,N
                                                                                ROS01830
   8
        V(I,J) = B(I,J)/R(J)
                                                                                ROS 01840
        RETURN
                                                                                ROS01850
        END
                                                                                RDS01860
C
                                                                                ROS 01870
       FUNCTION FUNXON(O,Z,N)
       VARIABLE X(I) IS TEMPORARILY SUBSTITUTED BY VARIABLE O(I)
                                                                                RDS 01880
C
                                                                                 ROS 01 890
       DIMENSION 0(40), Z(40)
```

FUNXON=100.\*(0(2)-0(1)\*0(1))\*\*2+(1.-0(1))\*\*2

RETUR'I Fi.D

С

ROSO1910 ROSO1920 ROSO1930 RUSO1940 ROSO1950

## APPENDIX D2

## CONFIDENCE INTERVAL CALCULATION FOR PARAMETER ESTIMATES

The confidence intervals for the parameters were calculated from the observations as follows:

- 1. The first order partial derivatives of the rate expression with respect to each of the parameters were obtained.
- 2. The numerical values of the derivatives were obtained by substituting the values of independent variables i.e. concentration and catalyst ratio and the estimated values of parameters. The values were written in the form of a nxm matrix. This nxm matrix is denoted by x where m being number of parameters and n being number of observations.

Let

$$R = \frac{b_1[RH]^2}{b_2 + b_3[M]}$$

be the rate expression.

Differentiating with respect to b,

$$\left(\frac{\partial R}{\partial b_1}\right) = \frac{\left[RH\right]^2}{b_2 + b_3 \left[M\right]}$$

Similarly, the other two derivatives are

$$\left(\frac{\partial R}{\partial b_2}\right) = \frac{-b_1 \left[RH\right]^2}{\left(b_2 + b_3 \left[M\right]\right)^2}$$

$$\left(\frac{\partial R}{\partial b_3}\right) = \frac{-b_1[RH]^2[M]}{\left(b_2 + b_3[M]\right)^2}$$

The matrix is of the form

$$X = \frac{(\frac{\partial R}{\partial b_1})_1}{(\frac{\partial R}{\partial b_2})_1} \frac{(\frac{\partial R}{\partial b_2})_1}{(\frac{\partial R}{\partial b_2})_n} \frac{(\frac{\partial R}{\partial b_3})_1}{(\frac{\partial R}{\partial b_3})_n}$$

Where the subscript indicates the number of experimental observation. For example, in case of cumene a 14 x 3 matrix was obtained as 14 was the number of experimental observations.

3.  $[X^TX]^{-1}$  was obtained by inverting  $[X^TX]$ .  $[X^TX]^{-1}$  was a 3 x 3 matrix in all the cases.

$$c = [X^{T}X]^{-1} = \begin{bmatrix} c_{11} & c_{12} & c_{13} \\ c_{21} & c_{22} & c_{23} \\ c_{31} & c_{32} & c_{33} \end{bmatrix}$$

A typical  $[X^TX]^{-1}$  for the system cumene was,

The parameter values for cumenc obtained by Rosenbrock optimization procedure are as

$$b = \begin{bmatrix} 2.2588 \times 10^{2} \\ 1.3744 \times 10^{10} \\ 1.2048 \times 10^{3} \\ 1.2$$

The individual parameter confidence intervals were calculated using the equations

where

 $s_{y_i}^-$  - estimated variance of y

 $t_{1-}$   $\frac{\alpha}{2}$  - value from 't' distribution corresponding to to a significant level of  $\alpha$  in a two sided test.

 $s_{\overline{y}_i}$  was obtained from the expression

$$s_r = \sqrt{s_r^2} = \sqrt{\frac{RSS}{\gamma}}$$

where RSS - residual sum of squares

 $\dot{\gamma}$  - degrees of freedom

= (number of observations - number of parameters).

APPENDIX D3

RESIDUAL ANALYSIS

TABLE D.1: SYSTEM-CUMENE

Observed Rate g mole/sec x 10 <sup>7</sup>	Predicted Rate g mole/sec x 10 <sup>7</sup>	Hydrocarbon concentration g mole/lit	Residual x 10 <sup>7</sup>
4.800	5.030	7.820	-0.230
4.130	3,220	6.260	0.910
1.730	1.810	4.690	-0.080
1.140	1.310	3 <b>.9</b> 90	-0.170
0.667	0.805	3.130	-0.138
4.380	5.030	7.820	-0.650
3.430	3.220	6.260	0.210
2.170	1.810	4.690	0.360
1.220	1.310	3.990	-0.090
0.667	0.805	3.130	-0.138
3.370	3.220	6.260	0.150
1.900	1.810	4.690	0.090
1.500	1.310	3.990	0.190
0.718	0.805	3.130	-0.089

TABLE D-2: SYSTEM-CYCLOHEXENE

Observed Rate g mole/sec x 107	Predicted Rate, g mole/sec x10 <sup>7</sup>	Hydrocarbon concentration g mole/lit	Residual x 10 <sup>7</sup>
4.90	4.89	7.88	0.0119
2.50	2.76	5.92	-0.2590
2.08	1.91	4.93	0.1670
1.50	1.23	3.94	0.2570

TABLE D-3: SYSTEM-TETRALIN

Observed Rate,7 g mole/sec x107	Predicted Rate, g mole/sec x 10 <sup>7</sup>	Hydrocarbon concentration g mole/lit	Residual x 10 <sup>7</sup>
12.00	12.00	7.37	0.0350
7.84	7.64	5.89	0.1960
3.86	4.30	4.42	-0.4440
3.00	2.99	3 <b>.</b> 68	0.0081